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PATENT SPECIFICATION

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COMPLETE-SPECIFICATION

Improved Process of High Vacuum Distillation

We. RASPIGLEY KOULK CONCLUST, a mm. Hg, and preferably less than 0.1 Company organised under the Lews of mm. Hg.) and to provide a purchase the State of New Jersey, United States of whereby such materials may be purised almerica, of 484, States Street, Rochester, without undesirable accomposition and States of New York, United States of with a considerable saving in time and America, Glasgiances of KENNETHE (ALDRE) heat onesylvation and the control of the saving in the same of the saving in the saving saving in the saving saving in the saving savin

America, (Assgrases of INSPIRERY ULAUDE DEVENBURY HUMEARY, a British Subject, of Kodak Paris, Ronhester, Oomity Monroe, State of New York, United States of 10 America), do hereby declars the nature of this invention and in what manner the same is to be performed, to be particularly to the present of the property of the particularly following statement:—of in and by the following statement:-

This invention relates to an improved process for the vacuum distillation of process or the vacuum unsurance or organic substances such as vegetable and animal oils, fats and waxes containing vitamins, sterols or hormones and which 20 are liquid at the temperature of distilla-

tion.

Processes of high vacuum distillation are known in which difficultly volatile materials are removed in pure form from 25 impurities and undesirable admixtures. Thus in U.S. petant to Burch , 1965, 231, difficultly volatile oils are vacuum distilled to remove desired constituents of the control of t

80 natural oils and fats are vacuum dis-tilled to concentrate compounds of therapeutic value contained therein, such as vitamins.

Such as vitamins.

In such high vacuum distillation pro85 cesses, due to the difficultly volatile
nature of the compound being purified,
temperatures considerably above the boiltemperatures considerably above the boli-ing point of the desired compound under the vacuum obtaining must be used. A satisfactory removal of the desired com-pound at temperatures lower than those point as temperatures lower than those heretofore found necessary is of considerable importance, since it would result in a material saving in material 45 otherwise lost due to decomposition. The amount of hest required to volatilize the desired material would also be considerated to the considerate of the considera

ably less. The present invention has for its object to overcome the deficiencies of object to overcome the deducates of hitherto known processes for concen-trating or purifying materials by disti-lation at high vacua, (not more than 1 heat energy. It has already been pro-posed to isolate the female sex hormone 60. by the high vacuum-shortpath distilla-tion of placenta dispersed in oil, i.e. a

tion of placenta dispersed in cil, i.e. a solid dispersed in a liquid. According to the present invention the organic substance, such as an cil, fat or 65 wax, which contains vitamins, sterols or hormones and which is liquid at the tem-perature of distillation is mixed with a perature of distillation is mixed with a compound or mixture of compounds his-ing a boiling point, under the distilla-tion conditions obtaining, in the neigh-bourhood of that of the material which it is desired to purify; the added material is then preferably removed from the mixed distillate.

mixed distillate. When the materials being concen-when the materials being concen-tually materials are distilled, it has been above the boling point of the desired above the boling point of the desired material must be used. Thus, in dis-lilling fish also such as cod liver cill, the vitamina distill over at temperatures with the control of the control of the control of the material must be used. Thus, in dis-bution of the control of the control of the vitamina distill over at temperatures. optimum temperature for recovery of strains A being about 180' O. Recently 85 nearly pure vitamin A has been secured and it has been found that it distills at less than 100' O. under molecular distillation conditions, i.e. those conditions less than AUT. Or more molecular custilistion conditions, i.e. those conditions to the condition to or the condition of the c

[Price 1/-]

contained therein, the addition of a quantity of other material having approximately the same boiling point under vacuum conditions as the desired 5 vitamin, makes it possible to readily dis-

till the vitamin, using a much lower tem-

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perature than previously possible.

The use of such lower temperatures is a very desirable feature where heat 10 sensitive and easily decomposable materials are to be recovered. Further more, many of the desired distillates are solid or so viscous as to collect in the still and are difficult to remove. The addi-

15 tion of a liquid material having a boiling point in the neighbourhood of the boiling point of the desired distillate will prevent such clogging and make continuous operation possible. Thus, the addition of fetty acids boiling in the neighbourhood of sterols being distilled from an

or may hood of sterols being distilled from an oil prevent the solid or viscous distillate from donging the apparatus with resultant delay for cleaning.

The materials to be added may be selected from widely different types of the control of the

material may be used as long as it has a boiling point in the neighbourhood of 80 the distillate desired, under molecular distillation conditions as hereinbefore defined and has no adverse effect on the material undergoing treatment. Thus, for example fatty acids, esters, mineral

for example faity soids, estars, mineral soil fractions, terpuese, seential oils, have been found to give useful results. Of course, a compound which is subject to material decomposition abould not be selected. Altiphatro phthalates, beary 40 digytoerol tetrapropionate boil below vitamin A and are useful seasts to employ in its recovery. Many of the added materials are so ex45 pensive that they cannot be used without careful recovery. The best of the selection of the course of the selection of the course of the selection of t

vitamin or other material distilled may be run through a separate still where the 50 vitamin or other desired material may be separated. The added material will remain as the still residue, if it had a higher bottling point than the desired purified material, or will be in the distillate, it has a lower beling point. In covered practically quantitatively. The added material may still contain small amounts of the desired purified material, and but as they are used repeatedly, being

60) but as they are used repeatedly, being added again and again to the fresh material initially treated, this does not represent a loss. Obviously other methods of separation known in the art, such as 65 freezing, crystallization, solvent extrac-

tion or purely chemical means may be employed instead of distillation for separating the added materials.

As previously indicated, the added material may have a boiling point the meserial may have a poining point the same as, above or below that of the sub-stance to be removed. Preferably an added material having a boiling point below that of the desired substance is used, since the lower boiling material generally gives botter results at lower emperatures.

As pointed out above, the invention is applicable to any process of high vacuum distillation in which a difficultly volatile distillation in which a difficultity volatile organic material is separated as the distillate. Thus, vitanine A, D and/or E may be separated by distillation from vegetable and animal offs containing them, such as oul, halfbut, etc. Hwer olls, menhaden, salmon, diegrafi, wordther offs, and the distillation of the distillation

Hormones of various types may be re-covered from oily concentrates by the

covered from oily concentrates by the present invention, as may also sterols and 90 high boiling hydrocarbons from mixtures containing them.

The distillation may be carried out in the manner described in U.S. patents Nos. 1,825,856 and 1,942,856. Such 9 molecular distillation processes are generally extracted out at less than 1,1 min. generally extracted the second of the seco

C. A characteristic of this type of un-tillation is that the vapours are con-densed upon a surface located at a dis-tant than about the mean free densed upon a surface located at a dis-tance of less than about the mean free path of the molecules of residual gas. Fressures of above 1. mm. may be used 105 and it has been found that volatile materials in fish oils may be distilled at pressures as high sal mm. Since a more committed removal of non-relative

constituents takes place at pressures 110 below .01 mm, we prefer to operate at these lower pressures. However, as the added materials used enable a rapid recovery of the desired materials, it is not necessary in many cases to operate under 115 molecular distillation conditions and pressures higher than those normally

used in such processes may be employed, in which case the process may be termed one of pseudo-molecular distillation. We will now describe our invention by way of example as applied to a fish oil containing vitamins A and D.

We have found in our researches that, although pure vitamin A is an alcohol, 125 the vitamin A occurring in fish oils and highly prized for its medicinal value is a mixture of esters with only a small pro-portion of the free alcohol. The alcohol boils at about 95° C, and the esters at 180 160° to 190° C. under a molecular vacuum. The vitamin D from these oils boils at around 140° C. When, how-

boils at around 140° C. When, however, it is desired to drive the vitamins
5 off from the parent cil, it is found that
temperatures of 160° C. and 180° to 215° C.

with the second of the D and 180° to 215° C.

withmins are, present in the notes of the to
10 quantities and partly because there are
only relatively axed at mounts of fatty
glyverides in the oil of boiling points
similar to the vitamins. The oil of comtention of the vitamins of the relative the comtention of the vitamins of the complex of the comtree safty acids, but the fatty glyverides
with 8 to 14 carbon atoms in the side
chain are complexious by their absence.

chain are conspicuous by their absence.

chain are conspicuous by their absence. We therefore add, for instance, to the 20 incoming oil four parts of giverol triperlargonates (tri-pelargonin) and we then find that distillation of a fraction rich in A and D occurs at a temperature of 160° C. The quantity of the fraction 51 is about 6 %.

The added material, e.g. tripelargonin, serves another important purpose, in that it condenses with the desired frontion of the condenses with the desired frontion 30 sway more readily from the solid condensing surface. Our researches have shown that the distillation characteristics of Vitamin D and cholesterol are very similar. The vitamin D when ordin-35 artly eliminated has so small a bulk that it takes a long time to drain from the con-

it takes a long time to drain from the con-

55 arily eliminated has so small a bulk that it takes a long time to drain from the condensing surface and often suffere decomposition on the way. With cile rich in a condensing surface and often suffere decomposition on the way. With cile rich in a condensing surface in a mass of crystals. In melting these down, the vitamin D is volatilized and partly destroyed. It will be appreciated that when triplaignoin or it could be appreciated that when triplaignoin of the condensate is increased, the crystallization of the condensate in increased, the crystallization in the condensate in increased, the crystallization in the condensate in increased, the crystallization in the condensate in the c

larly important in pseudo-molecular dis-tillation where the molecules are con-densed on a surface which is at a greater distance than the mean free path of the molecules.

molecules.
The herein described invention constitutes a simple, economical and highly states a simple, economical and highly moved by vacuum distillation of difficulty volatile organic compounds and partieu; leady of heat sensitive difficulty volatile compounds from mixtures containing them. An outstanding advantage of our process is the more rapid distillation at 80 lower temperatures than could hereinforce 30 lower temperatures than could hereinforce 30.

sower temperatures than could herestofore 80 be used and the possibility of distilling solids without clogging of the apparatus. Having now particularly described and ascertained the nature of our said invention and in what manner the same is to 85 be performed, we declare that what we

claim 18:—

1. In the process of distillation of organic substances, which are liquid at the temperature of distillation and which 90 contain vitamins, sterols or hormones, to separate a desired fraction at a high separate a desired fraction at a high vacuum of not more than 1 mm. Hg, the step which comprises adding to the organic material to be distilled a sub-stance of low volatility, which distils over as a mixed distillate with a desired

reaction.

2. The process as claimed in claim 1 applied specifically to those substances 100 which are liquid at or near atmospheric

which are injust at or near atmospheric temperature.

In which a neutral oil containing vitamins, sterolo or hormones is distilled 105 in presence of an added material which distils with the desired fraction, after which the added material is separated from the mixed distillation.

4. The process as claimed in claim 1 110 applied to the separation of vitamin, sterol or hormone concentrates from material containing one or more of these therapeutic substances, the distillation being performed under molecular condi- 115 tions as hereinbefore defined.

5. The process as claimed in claim 1 which comprises adding to a substance to be yacuum distilled a material having a

be vacuum distilled a material having a boiling point in the neighbourhood of 190 the desired fraction and subjecting the mixture to vacuum distillation.

6. The process as claimed in claim 1 which comprises adding to a natural oil a substance having a boiling point near 125 that of the desired fraction and subject-ing the mixture to distillation at a long the mixture with the subject-ing the mixture with the subject of the persister between 70 and 250° C. and con-densing the distillate at a distance from 180°.

the evaporative surface of less than the mean free path of the molecules of resi-dual gas.

7. The process as claimed in any of the preceding claims in which the added material is more volatile than the desired fraction.

8. The process as claimed in any of the preceding claims in which the distilla-

tion is performed at pressures less than 10 0.1 mm, Hg. 9. Processes of high vacuum distilla-tion, substantially as described.

Dated this 9th day of October, 1936. W. P. THOMPSON & CO., 12, Church Street, Liverpool 1, Chartered Patent Agents.

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